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# Paleobiology



Aragonite bias exhibits systematic spatial variation in the late Cretaceous Western Interior Seaway, North America

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# SCHOLARONE<sup>™</sup> Manuscripts

# 1 Aragonite bias exhibits systematic spatial variation in the late

# 2 Cretaceous Western Interior Seaway, North America.

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# 8 ABSTRACT

9 Preferential dissolution of the biogenic carbonate polymorph aragonite promotes 10 preservational bias in shelly marine faunas. Whilst field studies have documented the impact 11 of preferential aragonite dissolution on fossil molluscan diversity, its impact on regional and 12 global biodiversity metrics is debated. Epicontinental seas are especially prone to conditions 13 which both promote and inhibit preferential dissolution, which may result in spatially extensive 14 zones with variable preservation. Here we present a multi-faceted evaluation of aragonite 15 dissolution within the late Cretaceous Western Interior Seaway of North America. Occurrence 16 data of molluscs from two time intervals (Cenomanian-Turonian boundary, early Campanian) 17 are plotted on new high-resolution paleogeographies to assess aragonite preservation within 18 the seaway. Fossil occurrences, diversity estimates and sampling probabilities for calcitic and 19 aragonitic fauna were compared in zones defined by depth and distance from the seaway 20 margins. Apparent range sizes, which could be influenced by differential preservation potential 21 of aragonite between separate localities, were also compared. Our results are consistent with 22 exacerbated aragonite dissolution within specific depth zones for both time slices, with 23 aragonitic bivalves additionally showing a statistically significant decrease in range size 24 compared to calcitic fauna within carbonate-dominated Cenomanian-Turonian strata. 25 However, we are unable to conclusively show that aragonite dissolution impacted diversity 26 estimates. Therefore, whilst aragonite dissolution is likely to have affected the preservation of

fauna in specific localities, time averaging and instantaneous preservation events preserve regional biodiversity. Our results suggest that the spatial expression of taphonomic biases should be an important consideration for paleontologists working on paleobiogeographic problems.

31

32 Key words: Mollusca, calcite, OAE2, Cretaceous, fossil record bias, sampling bias.

33

# 34 INTRODUCTION

35 WHILST the fossil record provides a unique window into past life on Earth, it is well known 36 that it is both pervasively and non-uniformly biased (Raup, 1976; Koch, 1978; Foote and 37 Sepkoski, 1999; Alroy et al., 2001; Allison and Bottjer, 2011). Geologic, taphonomic and 38 anthropogenic biases (such as the amount of available fossiliferous rock for sampling, variation 39 in fossilization, and the degree to which the available rock record has been sampled) skew or 40 remove information from the fossil record, leaving the remaining catalogue of data uneven and 41 incomplete. Although biomineralized remains have an increased preservation potential 42 compared to soft bodied tissues (Allison, 1988; Briggs, 2003), they are still influenced by 43 various geologic and taphonomic processes (Kidwell and Bosence, 1991; Kidwell and 44 Brenchley, 1994; Kidwell and Jablonski, 1983; Best, 2008; Hendy, 2011). Shelly marine 45 faunas are especially susceptible to misrepresentation due to preferential dissolution of 46 biogenic carbonate polymorphs. It is well established that aragonite, a polymorph of CaCO<sub>3</sub> 47 found within the biomineralized shells of many invertebrates, dissolves more rapidly than the more common form of CaCO<sub>3</sub>, calcite, and at a higher pH (Canfield and Raiswell, 1991; Tynan 48 49 and Opdyke, 2011). Whilst both polymorphs can be destroyed by adverse conditions near the 50 sediment-water interface (Best and Kidwell, 2000) and the effects of dissolution can vary 51 between fauna (due to microstructure surface area, morphology, and shell organic content:

52 Walter and Morse, 1984; Harper, 2000; Kosnik et al., 2011), it is still the case that aragonitic 53 shells are more likely to dissolved than calcitic remains (Brett and Baird, 1986). As mineral 54 composition of molluscs is usually conserved at the Family level (Carter, 1990), this has the 55 potential to skew the record of molluscan diversity and trophic structure through time (Cherns 56 et al., 2011; Cherns and Wright, 2000; Wright et al., 2003), and negatively affect subsequent 57 work that relies on the relative abundance and distribution of shelly marine fauna (Kidwell, 58 2005). Cherns and Wright (2001) argued that early-stage dissolution could be substantial and 59 referred to the phenomenon as the "Missing Mollusc" bias. Subsequent work on a multitude 60 of temporal and spatial scales (Wright et al., 2003; Bush and Bambach, 2004; Kidwell, 2005; 61 Crampton et al., 2006; Valentine et al., 2006; Foote et al., 2015; Jordan et al., 2015, Hsieh et 62 al., 2019) has debated the magnitude of this bias; however, there is a broad agreement on the 63 potential for dominantly aragonitic shells to suffer greater post-mortem diagenetic destruction 64 in the Taphonomically Active Zone (TAZ) (Davies et al., 1989; Foote et al., 2015). Whilst the 65 effects of dissolution on the global macroevolutionary record of molluscs has been found to be 66 limited, possibly due to the potential of aragonite to recrystallize to calcite (Kidwell 2005; Paul 67 et al., 2008; Jordan et al., 2015), it is conceivable that local or regional conditions could impact 68 severely on perceived patterns of biodiversity in restricted areas (Bush and Bambach 2004). In 69 a regional study of Cenozoic molluscs, Foote et al. (2015) found evidence to suggest that 70 aragonite dissolution was both enhanced in carbonate sediments and insignificant within 71 siliciclastic sediments, with similar preservation potential of aragonitic and calcitic fauna 72 within the latter. They further emphasized the fact that scale is an important factor in determining the observable impacts of aragonite dissolution, which will strongly vary between 73 74 local (potentially consisting of an individual bed), regional and global studies. However, to 75 date research has focused on assessing the influence of aragonite bias on temporal trends of 76 biodiversity and has ignored the potential for direct spatial expression.

77 Early stage dissolution occurs within modern environments as a result of microbially mediated 78 reactions increasing local acidity (Walter et al., 1993; Ku et al., 1999; Sanders, 2003). 79 Bacterially-mediated decay of organic material within the upper sedimentary column occurs in 80 a series of preferential redox reactions. By-products of these reactions, such as solid phase sulphides from sulphate reduction and CO<sub>2</sub> from aerobic oxidation, result in changes to local 81 82 pore-water saturation of calcium carbonate (Canfield and Raiswell, 1991; Ku et al., 1999). 83 Additionally, oxidation of H<sub>2</sub>S above the oxycline increases acidity at that boundary; if this 84 occurs at the sediment-water interface then it can adversely affect the preservation of shelly 85 marine fauna (Ku et al., 1999). As such, dysoxic sedimentary environments might have a 86 predisposition for dissolution of biogenic carbonate and enhance the effect of the "Missing 87 Mollusc" bias (Jordan et al., 2015). Epicontinental seas, marine water-bodies which form by the flooding of continental interiors, are especially prone to strong water column stratification 88 89 and sea level variation, and have a pre-disposition to seasonally anoxic or dysoxic conditions 90 (Allison and Wells, 2006; Peters, 2009). As such, they have the potential to be more prone to 91 both preferential aragonite loss and preservation than modern oceans. Cherns et al.'s (2011) 92 model for taphonomic gradients of aragonite preservation along a shelf to basin transect can be 93 readily applied to epicontinental sea settings (Fig. 1). If we assume the center of a seaway was 94 stratified with at least a seasonally anoxic basin floor, we would expect enhanced dissolution 95 to occur in the seaway margins, likely in the mid-to-outer shelf setting (Cherns et al., 2011). In 96 the anoxic basin center we would expect to see enhanced preservation, as an aragonitic skeleton 97 residing on the surface sediment in an anoxic water column would not be susceptible to 98 dissolution from H<sub>2</sub>S oxidation (Jordan et al., 2015; however, we would not expect to see 99 abundant benthos in such a setting because of bottom water toxicity). It is apparent this could 100 result in spatially expansive zones with conditions predisposed for heightened aragonite 101 dissolution and preservation (Fig. 1; it is important to note that we do not expect all aragonitic

fauna to be missing from any region of the seaway – merely that a lower relative proportion of aragonitic molluses be found, due to a reduced probability of an individual site recording their occurrence). How these hypothesized basin-margin to basin-center zones could influence long term patterns of molluse distribution, preservation and recovery remains to be examined. As epicontinental seas contain the majority of our Phanerozoic fossil record (Allison and Wells, 2006), it is imperative that we understand systematic biases that may specifically affect these settings.

109

Here we present a spatial investigation of aragonite dissolution within the late Cretaceous Western Interior Seaway (WIS) of North America, using sampling probability estimates and multiple logisitic regression to evaluate patterns of spatial distribution in preserved calcitic and aragonitic fauna. We address two key questions: (1) does aragonite bias exhibit systematic spatial variation across the seaway and (2) if so, does this influence perceived patterns of diversity?

116

# 117 MATERIALS AND METHODS

118 *Time Intervals and Paleogeography* 

The two stratigraphic intervals or time slices (Cenomanian-Turonian and early Campanian) were selected: (1) because of purported dysoxic conditions within their duration; and (2) due to their differences in environment, oceanography and preserved lithology, allowing for comparison of taphonomic regimes. The first interval covers the Cenomanian–Turonian boundary, spanning from the *Dunveganoceras pondi* to *Collignoniceras woollgari* ammonite zone (~94.7–93 Ma) (Cobban et al., 2006). The second interval spans the early Campanian, from the *Scaphites leei III* to *Baculites obtusus* ammonite zones (~83.5-80.58 Ma) (Cobban et

126 al., 2006). The geologic context of stratigraphic intervals is detailed in Supplementary127 Information 1.

128 A global atlas of 1:20,000,000 scale paleogeographic maps, compiled by GETECH plc, formed 129 the basis for new regional-scale, high resolution interpretations for the selected time intervals. 130 The original paleogeographic maps (Markwick, 2007) are underpinned by the GETECH plate 131 model (v1), which is outlined further in Supplementary Information 1. High resolution 132 mapping involved synthesis of stratigraphic, sedimentologic and paleontologic information to 133 produce 1:5,000,000 scale paleogeographies with suggested paleobathymetry. A full list of 134 decisions on paleogeographic reconstructions and key references for each time interval are 135 provided in Supplementary Information 1.

136 Landward-to-basinward arrangements of *a priori* binned zones for each time-slice were based 137 on average paleobathymetry (Fig. 2). Bathymetric reconstructions were divided into four bins, 138 each of which covers a specific interpreted depth range: Nearshore (<50 m), Proximal Offshore 139 (50-100 m), Distal Offshore (100-150 m) and Basin Center (>150 m). These designations were 140 based on the previously constructed paleobathymetry for the WIS produced by Sageman and 141 Arthur (1994), but match the paleobathymetry in our new maps and represent a reasonably 142 high resolution without being compromised by large changes in shoreline position within our chosen time slices. 143

Distance-from-paleoshoreline zones (Fig. S1) were constructed based on 50 km intervals from the time-averaged paleo-shoreline position until reaching the basin center, with number of occurrences, collections and total outcrop area plotted per zone. These were generated by constructing a fishnet of points in ArcGIS (ESRI, 2010) using the "Fishnet tool", which were selected by the "Select By Location" tool with increasing distance in 50 km intervals from the paleoshoreline: the position of the most basinward selected points was used for the bin

boundary. Results for depth zones are used in the main body of this manuscript; distance-frompaleoshoreline zones are available in Supplementary Information 1 and Figures S2, S4, S6 and
S7.

153

154 Fossil Dataset

155 A presence-only fossil occurrence dataset of bivalve and ammonite taxa was produced for the 156 selected stratigraphic intervals, collated from personally provided digitized collections from 157 the United States Geological Survey (USGS) and Smithsonian Museum of Natural History 158 (NMNH), as well as downloads from the Paleobiology Database (PBDB; http:// 159 paleobiodb.org), and iDigBio (http://www.idigbio.org). Each occurrence includes taxonomic 160 and geographic locality data, an associated collection with lithologic and geologic information, 161 and modern latitudinal and longitudinal co-ordinates. Data were extensively screened for problematic records and to ensure taxonomic validation (see Supplementary Information 2 for 162 163 the latter).

164 The resultant Cenomanian-Turonian dataset contains 5867 occurrences from 2409 localities, 165 with 207 genera, 1549 species, and 3886 specimens identifiable to species level. The early 166 Campanian dataset comprises 2544 occurrences from 1186 localities, recording 163 genera, 1405 species, and 1405 specimens identifiable to species level. Generic level taxonomic 167 168 diversity was used for all tests; species level results can be found in Supplementary Information 169 1 and in Supplementary figures S3-6. Full information regarding downloads, sources and 170 screening of data can be found in Supplementary Information 1, and the full dataset found in 171 Supplementary Information 2.

172

173 Mineralogy

174 Bivalve shells are a composite of layered mineral crystallites, which are sheathed by a 175 refractory organic matrix of fibrous protein (Taylor, 1969). As these mineral layers can be 176 comprised of both calcite and aragonite, variation in overall mineral composition must be taken 177 into account when assigning a predominant mineralogy to a specific bivalve Genera. Different 178 scoring mechanisms have been adopted by previous workers to address this issue. Kidwell 179 (2005) used a five-point decimal scoring system from entirely aragonitic (1) to entirely calcitic 180 (3), with three permutations of mineralogy between. Crampton et al. (2006) adopted a simple 181 and effective system of counting organisms as calcitic if they contained a calcitic element that 182 would allow them to be identified to species level. We utilise a combination of these 183 approaches - organisms were scored using the system of Kidwell (2005) to maintain the 184 maximum amount of data, but simplified into binary categories afterwards based on whether 185 they contained sufficient calcitic parts to enhance preservation potential. Note that we have not included either the inner myostracal layer or periostracum in our assignments of mineralogy. 186

187 Information on shell composition was predominantly gathered from a personally provided dataset from S. Kidwell (Kidwell, 2005), as well as further studies from Taylor (Taylor, 1969; 188 189 Taylor and Layman, 1972), Majewske (1974), Carter (1990), Schneider and Carter (2001), 190 Lockwood (2003), Hollis (2008) and Ros-Franch (2009), as well as many papers focussed on 191 single genera or families. For genera for which information regarding shell mineralogy was not 192 available, composition was assigned based on the dominant mineralogy of the family, as 193 composition is highly conservative both amongst species within a genera and amongst genera 194 within a family (Taylor, 1969). In total, 124 bivalve genera were assigned a mineralogy, of 195 which 41 (33%) were achieved using familial relation (Supplementary Information 2).

196

197 Life habits

Life habits of bivalves were assembled to allow additional interrogation and interpretation of environmental and sampling regimes. Life habits were separated into the following categories: relation to substrate, mobility and diet. Data for each genera of bivalve were primarily gathered from the NMiTA Molluscan Life Habits Database (Todd, 2017) and the PBDB, with further data collected from the wider literature (Supplementary Information 2).

203

204 Outcrop Area

205 Relevant rock outcrop area was plotted per zone to evaluate broader scale bias influencing 206 patterns of fossil distribution. Outcrop areas for the selected time slices were generated by 207 combining state-wide digitized geologic maps from publicly available USGS downloads and 208 selecting shape files which matched formations found within those time slices. Some State 209 Surveys grouped relevant formations with other partially contemporaneous formations that 210 spanned multiple stages: we chose to include these designations in order to present the 211 maximum possible sampling extent in terms of outcrop area. Outcrop was projected in ArcGIS 212 (ESRI, 2010) using the USA Contiguous Albers Equal Area conic projection, to minimize 213 distortion of distances. Outcrop areas per zone were created by using the "Intersect tool" in the Geoprocessing toolbar in ArcGIS, and area (km<sup>2</sup>) calculated using the Calculate Geometry 214 function in the attribute table. Outcrop was split into depth zones by using the Intersect tool in 215 216 ArcGIS (ESRI, 2010). Outcrop area for each zone was calculated by summing the total area of 217 all outcrop polygons within that zone. Collections per zone were counted by exporting 218 occurrences selected in zones in the seaway as shapefiles, then using the "arcgisbinding" 219 package to view and organise the data in R version 3.0.2 (Team, 2017).

220

221 Dominant lithology

Each collection was assigned a dominant lithology to allow for comparative testing. If these data were not available, a lithology was assigned from the dominant lithology of the formation, with reference to USGS formation records. Collections were assigned one of the following lithologies (primarily based off original USGS records): siliciclastic mudstone, siliciclastic siltstone, siliciclastic sandstone, conglomerate, ironstone, calcareous mudstone and siltstone, marl, calcarenite, limestone and chalk.

228

229 Range Size

230 If the presence of preferentially destructive zones is affecting the spatial distribution of 231 aragonitic fauna, we might expect to see overall smaller range sizes for aragonitic organisms 232 compared to calcitic organisms (Fig. 3). As such, range size estimates were produced for 233 calcitic and aragonitic bivalves and compared to test if aragonite bias influenced perceived 234 range of aragonitic organisms. Note that ammonites were excluded from this test due to the 235 difference in life habit between them and bivalve fauna: ammonites have a pelagic to nektono-236 benthic mode of life (Ritterbush et al., 2014), whilst bivalves are predominantly epifaunal and 237 infaunal.

238 Geographic locality data for the selected fauna was visualized in ArcGIS (ESRI, 2010). Faunal 239 occurrences were paleo-rotated using the Getech Plate Model to match the paleogeography of 240 the appropriate stages of the Late Cretaceous. This ensures that tectonic expansion and 241 contraction of the North American plate from the Mesozoic to Recent has a negligible effect 242 on propagating estimation error in range-size reconstructions. Fossil occurrences were projected into ArcGIS using the using the USA Contiguous Albers Equal Area conic 243 244 projection. A 10 km buffer was additionally applied to each occurrence point in order to control 245 for any error in paleogeographic or present position of fauna. ArcGIS (ESRI, 2010) was then 246 used to construct convex hull polygons for each taxon, and the spatial analyst tools from this

247 software calculated the area of each reconstructed polygon. We did not account for landforms 248 within the ranges of any organisms, and thus ignored their area when calculating overall area 249 of ranges. Several vertices for range size polygons appeared on what is classified as land within 250 our paleogeographies; due to rapid changes in shoreline position within the WIS, we decided 251 to keep using these fauna for range size estimations. Myers and Lieberman (2011) showed that 252 relative range sizes for vertebrates in the WIS were not overly affected by resampling 253 occurrence points - consequently, we have not carried out a similar test for this study. 254 Comparisons between the ranges of aragonitic and calcitic fauna were carried out using the 255 Wilcoxon-Mann-Whitney test with continuity correction (Brown and Rothery, 1993). 256 Geographic range data for all applicable taxa are provided in Supplementary Information 2.

257

# 258 Sampling Probability and Multiple Logistic Regression

To be able to further observe differences between aragonitic and calcitic organisms throughout 259 260 the seaway, we employed a modified version of the sampling probability method used by Foote 261 et al. (2015) (after Foote and Raup, 1996). In this method, the sampling probability of a time 262 bin was generated by compiling a list of all fauna with originations older than that bin and extinctions younger, and then dividing the total number of species found within the bin by that 263 264 figure. This allows for a sampling probability to be estimated on a per bin, per group basis. 265 Here we devised three variants on this method for application in the spatial realm. It should be 266 made clear that the modified methods utilized in this work come with the caveat that in the 267 spatial realm it is impossible to know if a species was present in a precise location in the past: for instance, if zones A, B, and C are designated with increasing distance away from a 268 269 paleoshoreline, it cannot be assumed that because an organism exists in zones A and C that it 270 was ever present in zone B. Consequently, the probabilities generated from the methods 271 described below are relative, and cannot be taken as a "true" probability. However, the methods

272 utilized were designed to be as inclusive as possible and to deliver a strongly conservative 273 estimate of true sampling probabilities between groups; consequently, these methods provide 274 a useful estimate on the relative likelihood of sampling aragonitic or calcitic fauna. 275 Furthermore, sampling probabilities through time based on regional studies such as those 276 utilized by Foote at al. (2015) rely on the assumption that groups were not genuinely absent 277 from the study region at a particular time and that other geographic variables do not have an 278 effect – as such the use of these metrics to evaluate the distribution of fauna across the WIS is 279 validated.

280 Three methods were devised for dealing with the issue of unknown "correct" distribution of 281 species across the seaway and to correct for differences in the number of collections between 282 zones: (1) finds two bins either side of the current bin and generates a list of the total number 283 of possible species across those five bins; (2) finds all formations that appear in the selected bin that contain specimens of the selected group (e.g. calcitic bivalves), and then finds the total 284 285 number of species for that group from those formations; (3) finds all formations in the current 286 bin and the two adjacent bins that contain specimens of the selected group, and subsequently 287 finds the total number of species from those formations. For all three methods, the total number 288 of sampling opportunities per bin was generated by multiplying the number of potentially 289 recoverable species by the number of collections to standardize for differences in collecting 290 intensity. The low number of depth-based bins could potentially result in flattening the curve 291 of sampling probability using Method 3, and thus Method 2 is employed in the main body of 292 this paper for depth-based results.

To determine the primary controls on sampling probability between the two stages, we used multiple logistic regression, coding sampling opportunities as the response variable and mineralogy, lithology, life habits (mobility, relation to substrate and feeding style) and depth zone as the predictor variables. Multiple logistic regression allows for the use of binomial

nominal values by using the odds ratio, a measure of the relationship between the odds of an outcome, in this case sampled (1) or not sampled (0), along with multiple potentially explanatory ecological or physiographic variables. A full model is generated that incorporates all potential variables, and a null model defined that includes none. Stepwise addition or deletion from the null or full models, respectively, and analysis in the change of likelihood and of respective AIC (Akaike information criterion) scores contributes to a final predictive model of explanatory variables and respective statistical significance.

Sampling opportunities were tabulated as the presence or absence of each recoverable genera per collection, per depth zone. Each sampling opportunity was assigned a lithology based on collection lithology, as well as all ecological attributes related to that genus. To test for multicollinearity between variables, correlation tests were run using Spearman-Rank correlation using the Performance Analytics package in R. Explanatory variables that showed a strong (above 0.7) statistically significant correlation were excluded from further analysis (Supplementary Information 1).

311 Interaction terms were also added to explore the possibility of multiple confounding factors 312 and increased model complexity. These terms were restricted to a combination of lithology and 313 mineralogy, so as to test for specific interactions between the two (e.g. whether preservation of 314 aragonite was specifically enhanced within limestones). We also partitioned the data to be able 315 to fully explore the influence of various contributing factors on sampling probability per depth 316 zone, as well as include all organisms in the data (ammonites were excluded from analyses 317 involving life habits, as discussed below). Both effect sizes of individual factors and AIC 318 values of models are presented for statistically significant interactions. All methods were 319 written and implemented using R.

320

321 Occurrences, Raw diversity and SQS

322 To establish the potential influence of aragonite bias on diversity of shelly taxa, total 323 occurrences of organisms were counted per zone using the Select By Location tool in ArcGIS 324 (ESRI, 2010) which were used to generate landward-to-basinward profiles of raw occurrences, 325 raw and subsampled diversity estimates. Shareholder quorum subsampling (SQS; Alroy, 326 2010), a method for standardising taxonomic occurrence lists based on an estimate of coverage, 327 was implemented in R using script provided by Alroy (pers. comms.) for each faunal group. 328 Calcitic and aragonitic groups were evaluated for statistically significant differences using the 329 Chi-squared test for non-random association (Brown and Rothery, 1993). All statistical tests 330 were implemented in R. Results pertaining to patterns within raw occurrences can be found 331 within Supplementary Information 1 and Figure S1.

- 332
- 333 **RESULTS**
- 334 **Sampling Probability**
- 335 Cenomanian-Turonian

3 336 For generic level sampling probability (Fig. 4a), aragonitic bivalves and ammonites show a similar trend for the first three depth zones. After this, sampling probability drops to 0 for 337 338 aragonitic bivalves (as none were recovered), whilst it increases to a peak for ammonites. 339 Calcitic bivalves record a higher sampling probability than ammonites or aragonitic bivalves 340 in all zones and show a basinwards increase in sampling probability.

- 341
- 342 Campanian

343 In the lower Campanian (Fig. 4b) ammonites have the highest sampling probabilities, showing 344 a level trend across the seaway with a pronounced trough in the distal offshore. Aragonitic 345 bivalves record a relative high sampling probability in the nearshore, followed by a sharp 346 decline for both proximal and distal offshore zones and an increase towards the basin center.

Calcitic fauna have a consistently higher sampling probability than aragonitic bivalves, but
lower than ammonites; they also show a level trend across the seaway, experiencing a peak in
the distal offshore.

350

# 351 Sampling probability between lithologies

352 Cenomanian-Turonian

353 For the Cenomanian-Turonian (Fig. 4c,e,g), ammonites show the same trends and relatively 354 little difference in absolute values between carbonate and siliciclastic sampling opportunities; 355 the greatest difference appears in the basin center, where sampling probability is higher in 356 carbonates. Aragonitic bivalves show a much larger difference, with siliciclastic opportunities 357 scoring consistently higher than carbonate opportunities, even during the large decline within 358 the proximal offshore. Calcitic bivalves show virtually no difference in sampling probability 359 until the basin center, where sampling probability within carbonate sampling opportunities 360 increases substantially.

361

362 *Campanian* 

For the Campanian (Fig. 4d,f,h), siliciclastic opportunities of ammonites score higher than carbonate except for within the nearshore. Aragonitic bivalves are not sampled within carbonate collections in either the nearshore, distal offshore or basin center; their sampling probability curve is virtually entirely made by appearances in siliciclastic sampling opportunities. Calcitic bivalves show a decoupled trend between lithologies, with carbonate sampling opportunities showing higher on average sampling probabilities that increase towards the basin center, compared to the fairly low scoring, level trend in siliciclastic.

370

# 371 Multiple Logistic Regression

Results of the logistic regressions are shown in Tables 1-8 and summarized in Fig 5. When interpreting these, note that calcitic mineralogy is compared to aragonitic, so that positive regression coefficients indicate greater odds of sampling calcite. As lithology has multiple parameters, these were compared against the relative sampling probability of mudstone, which is used as a baseline. We are primarily interested in reporting effect sizes, which are gauged by the magnitude of regression coefficients.

AIC scores are utilized in choosing ideal model fit when comparing models with and without two-way interactive terms (a combination of effects between explanatory parameters: for example, the relative odds of sampling calcitic fauna within a specific lithology), with lower scores indicating a better model fit. Only models with the lowest AIC scores are presented here and we only report factors with statistically significant results (p<0.05); full results can be found within Supplementary Information 2.

384

# 385 Cenomanian-Turonian

386 Mineralogy, lithology, feeding style and depth zone all influence the preservation potential of fauna in the seaway (Table 1); lower AIC scores when an interactive term is added suggest this 387 388 provides a better model fit than when this is excluded. The odds of sampling calcitic fauna are 389 shown to be 4.6 times (the exponential of the coefficient; 1.52) higher that of aragonitic fauna, 390 with ANOVA results showing mineralogy contributing the most towards deviance from the 391 null model. Limestone environments are shown to be detrimental to the sampling probability 392 of fauna, whereas sandstones and siltstone enhance sampling probability. The positive 393 interaction between mineralogy and limestone lithologies shows that aragonitic fauna have 394 comparatively strongly reduced odds of being sampled within limestone environments. All 395 depth zones are shown to have decreased sampling probability compared to the basin center, 396 with nearshore and proximal offshore zones showing the worst sampling potential.

397 Chemosymbiont deposit feeders are shown to have an increased preservation potential398 compared to other feeding styles.

399

400 We additionally partitioned the data into each depth zone, to test for differences with increased bathymetry across the seaway (Table 2). The nearshore zone exhibits an increase in the odds 401 402 of sampling calcitic fauna, although this effect is reduced compared to results across the whole 403 seaway. Sandstones are also shown to exhibit increased sampling probability. The proximal 404 offshore shows a significant increase in the odds of sampling calcitic bivalves relative to 405 aragonitic bivalves (6.17 compared to 1.88 for the nearshore), as well as increased sampling 406 probability in marl depositional environments and for chemosymbiotic deposit feeders. 407 Limestone negatively impacts the sampling probability of bivalves; the positive interaction 408 between calcite and limestone consequently suggests that this negative impact is related to the 409 sampling probability of aragonitic bivalves. The distal offshore shows a similar pattern, 410 although the relative odds of each are reduced compared to the proximal offshore. The basin 411 center shows increased odds of sampling bivalves within calcarenite, calcareous shale and marl 412 environments, but no other statistically significant terms.

413

We also assessed depth zones for the inclusion of all organisms (Table 3). When ammonites ae included, the odds of sampling aragonitic fauna increase (calcitic bivalves show odds of 2.1 higher sampling probability). Sandstone shows reduced odds of sampling any fauna, the opposite of previous results. The interaction between mineralogy and lithology shows increased sampling probability of calcitic organisms within limestones, sandstones, calcarenites and calcareous mudstones, suggesting this effect is predominantly produced by the addition of ammonite fauna.

421 When assessing zones independently (Table 4), nearshore sampling probabilities are only 422 controlled by mineralogy, although again with lower odds than reported elsewhere (1.56). In 423 the proximal offshore, results show an increased sampling probability of calcitic fauna within sandstones and calcareous mudstones. The distal offshore also shows strong interactions 424 425 between sampling probability of calcitic fauna and lithology, with strongly positive coefficients 426 for sandstone, limestone, calcareous shale, and calcarenite two-way interactions. Overall, the 427 sampling probability of calcite compared to aragonitic fauna is high, although reduced 428 compared to the proximal offshore. Within the basin center, mineralogy is not listed as a 429 statistically significant interactive term on its own, but calcitic fauna exhibit increased 430 sampling probability for interactive terms with calcarenites, calcareous mudstones, limestones, 431 and marls.

432

# 433 Campanian

Models for all bivalves in the Campanian (Table 5) show comparatively few statistically significant contributors to sampling probability. By comparison with the Cenomanian, bivalve samples from the Campanian are only weakly influenced by mineralogy (showing odds of 2.16 increased likelihood of sampling calcitic organisms). Additionally, only sandstone and interactions between sandstone and limestone with calcitic organisms are shown to exert any other influence on sampling probability.

This trend continues when partitioning the bivalve data into depth zones (Table 6). The nearshore zone has no statistically significant individual factors contributing to sampling probability. The proximal offshore includes statistically significant effects due to mineralogy and lithology, particularly limestones and calcareous mudstones where sampling probability is enhanced. Mineralogy, sandstone and the interaction between mineralogy and sandstone are reported as statistically significant factors for the distal offshore; mineralogy has a relatively

446 high positive coefficient (odds of 3.16 in favour of calcitic organisms). Sampling probability 447 is enhanced in sandstones overall, but negatively influences the odds of recovering calcitic 448 organisms: it therefore follows that aragonitic bivalves show particularly enhanced sampling 449 within sandstones. Model results for the basin center suggest that only calcareous shale has a 450 statistically significant positive impact on sampling probability.

451 When all organisms are assessed (Table 7), mineralogy and depth zone are the only 452 contributors to the full model. Surprisingly, aragonitic organisms have a higher sampling 453 probability than calcitic using the full model, with mineralogy only contributing to a very small 454 amount of deviance from the null ANOVA model. As this result is not observed when assessing 455 bivalve fauna, it is likely that ammonite occurrences are principally contributing to this effect. 456 Depth zones were also evaluated for all organisms (Table 8). Only the proximal offshore 457 supported a model other than the null, which reported mineralogy as a contributing factor; 458 unusually, calcitic fauna are shown to have a reduced sampling probability compared to Revie aragonitic. 459

460

461 **Range Size** 

462 Cenomanian-Turonian

463 Box plots were generated on a log scale to show differences in mean range sizes between 464 calcitic and aragonitic organisms (Fig. 6a). There is a visible difference in variability of range 465 size between groupings, with calcitic fauna showing an average larger range than aragonitic. 466 The Wilcoxon Mann Whitney test also showed a statistically significant difference between 467 the range sizes for the two groups (p value = 0.00405), with a reported difference in median 468 range size of 48,694 km<sup>2</sup>. As sample size varied between the groups, resampling measures were 469 carried out to test the accuracy of these results. A randomized bootstrap with replacement 470 calculating the difference between the means of range sizes was implemented 10,000 times in 471 R (Fig. 6c). Our recorded difference in the mean was shown to have an associated p value of
472 0.0172, showing statistical significance.

473

# 474 Campanian

Box plots were generated to show differences in mean range sizes between early Campanian calcitic and aragonitic organisms (Fig. 6b). Calcitic bivalves show higher variability in mean range size than aragonitic bivalves. However, the Wilcoxon Mann Whitney test showed no statistically significant difference between the two groupings (p value = 0.504) with a recorded difference in median range size of 13,540 km<sup>2</sup>, and a randomized bootstrap (Fig. 6d) with replacement recovered an associated p value of 0.1527 (non-statistically significant).

481

# 482 Raw Diversity and SQS

483 Cenomanian-Turonian

Within the Cenomanian-Turonian, broadly similar patterns of diversity occur in all groups (Fig. 7a,c) – peak diversity is within the distal offshore, with lowest values in the nearshore and basin center. Calcitic bivalves show proportionally enhanced diversity in the proximal offshore compared to the other faunal groups. These patterns closely align with the number of collections within each zone, but show limited similarity to zoned outcrop area.

Subsampled ammonite and calcitic bivalve diversity show a broadly similar pattern to their raw taxic diversity signals (Fig. 8a,e). The record of aragonitic bivalves (Fig. 8c) is too poor to resolve subsampled diversity for the basin center; however, a slight decline in subsampled generic richness exists in the proximal offshore.

493

494 Campanian

495 Calcitic bivalves and ammonites exhibit a similar pattern in diversity (Fig. 7b,d) although the 496 latter show an increase in the proximal offshore. Aragonitic bivalve diversity has a similar peak 497 in the proximal offshore but declines towards the basin center. None of these trends show 498 similarity to the distribution of collections or outcrop area throughout the seaway.

When subsampled (Fig. 8b,d,f), calcitic and aragonitic bivalves are most diverse within the
proximal offshore, falling to relative lows within the distal offshore and basin center.
Ammonites are most diverse in the nearshore, followed by a decline to a flat profile.

502

# 503 **DISCUSSION**

# 504 Sampling probability and multiple logistic regression

505 Our results from estimations of sampling probability and subsequent multiple logistic 506 regression suggest that aragonite bias may be present within distinct depth zones of the seaway 507 during the Cenomanian-Turonian. Mineralogy has a strong and statistically significant impact 508 on sampling probability within the proximal and distal offshore bathymetric zones, and shows 509 the highest contribution to deviance from the null model. This is further supported by the fact 510 that whilst all aragonitic taxa have lower sampling proportions overall, both aragonitic bivalves 511 and ammonites disproportionally decrease in sampling probability within the proximal 512 offshore compared to calcitic bivalves. Ammonites, whilst still showing reduced sampling 513 probability compared to calcitic fauna, are more likely to be sampled than aragonitic bivalves; 514 a potential explanation for this difference could be that aragonite dissolution acts differently 515 upon ammonites compared to bivalves. Body sizes of ammonites and bivalves differ, with 516 ammonites generally having larger forms (Jablonski, 1996). This has been known to influence 517 preservation potential and the extent of aragonite dissolution: Wright et al. (2003) showed that 518 ammonites are affected less severely than aragonitic bivalves by early stage aragonite

519 dissolution, often exhibiting poor preservation rather than complete removal. Our results have520 the potential to be partially related to this effect.

521 Aragonitic bivalves have lower absolute sampling probabilities in carbonate environments than 522 in siliciclastic environments, supporting the results of Foote et al. (2015). However, when 523 examining the proximal offshore zone, we can see that sampling probability within siliciclastic 524 lithologies falls dramatically. As this zone records the largest difference in odds of sampling 525 between calcitic and aragonitic taxa, it can be argued that aragonite bias can influence fauna 526 within siliciclastic deposits in epicontinental seas, in contradiction to Foote et al. (2015). The 527 absolute sampling proportions of calcitic bivalves remain relatively consistent (at about 2% of 528 genera per collection) throughout the seaway until the basin center, where they increase 529 dramatically within carbonates compared to siliciclastics. Foote et al. (2015) reported that 530 calcitic organisms experienced higher sampling probabilities in carbonate-rich intervals, which is especially enhanced in limestones. As carbonates make up 93% of total sampling 531 532 opportunities within this zone, our results align fairly closely with previous findings. Whilst 533 Foote et al. (2015) singled out lithology as an important factor for aragonite dissolution, they 534 did not investigate whether differences in grain size significantly influenced results. Within 535 this study, sandstone and siltstone are consistently shown to have better odds at preserving 536 aragonitic fauna than mudstone. This is unsurprising, considering that coarser, oxidized 537 sediments are likely to contain lower quantities of organic matter than finer sediments, and thus 538 provide less material for the microbial decay which ultimately controls the dissolution of 539 aragonite within the taphonomically active zone (Cherns et al., 2008). However, siltstone 540 appears to have higher odds than sandstone, potentially a reflection of increased quality of 541 preservation in lower energy settings. It should be noted however that only a few models 542 include both siltstone and sandstone and therefore allow for comparison of sampling probabilities. 543

544 Potential ecological signals can also be parsed from the results reported here. Within the 545 Cenomanian-Turonian dataset, odds of sampling chemosymbiont deposit feeders within the 546 proximal offshore were higher than for other bivalves, forming a statistically significant part 547 of the final model and accounting for the second highest deviance from the null model. 548 Chemosymbiosis in bivalves occurs in a range of environments to cope with life in sulphide-549 rich environments, typically at deep sea vents or in sediments at the oxic/anoxic interface 550 (Cavanaugh, 1994). Combined with evidence for poor sampling probability of aragonitic fauna 551 in siliciclastic lithologies, this lends credence to the likelihood of fluctuating benthic oxygen 552 conditions within the proximal offshore, ideal for preferential aragonite dissolution. More 553 broadly, several previous works have suggested that aragonite bias strongly influences 554 perceived trophic communities within molluscan fauna, favouring preservation of specific life 555 habits (Cherns et al., 2008; Cherns and Wright, 2009). Unfortunately, very few statistically 556 significant life habit factors contribute to our final models (Fig. 5), and thus we cannot draw 557 any conclusions regarding preservational shifts in trophic structure. In the basin center, 558 ammonites are more likely to be sampled compared to other organisms. This confirms 559 expectations of enhanced preservation within a predominantly anoxic water column, where 560 dissolution and predation have reduced impact on the removal of fauna emplaced by pelagic 561 fallout (Jordan et al., 2015).

Within the Campanian, there is a somewhat contradictory pattern. Multiple logistic regression results show that mineralogy only has a strong, statistically significant impact on relative sampling odds when assessing bivalves within the proximal and distal offshore bathymetric zones, with only the latter showing a strong deviation from the null model in ANOVA results. When ammonites are added, the odds of sampling aragonitic fauna are actually higher than that of calcitic organisms within the proximal offshore, and all other zones show no statistically significant contributions from mineralogy. This is reinforced when one considers the absolute

569 proportions of mineralogies sampled: ammonites exhibit the highest overall sampling 570 probability between fauna. A potential cause of this contradiction is preferential sampling bias. 571 Ease of collecting and human interest can result in skewed sampling effort and intensity, potentially inflating (Foote and Sepkoski, 1999) or reducing (Lloyd and Friedman, 2013) the 572 published records of certain taxa, locations, and time periods above others. The WIS has long 573 574 been known for its abundance and diversity of ammonite fauna, and consequently ammonites 575 have been used for systematic biostratigraphic correlation since the 1930s (Stephenson and 576 Reeside Jr., 1938). An intensive effort to collect ammonites for stratigraphic purposes was carried out by a selection of workers through the latter half of the 20<sup>th</sup> century to the present 577 578 day (Scott and Cobban, 1959; Gill and Cobban, 1973; Cobban and Hook, 1984; Cobban et al., 579 2006; Merewether et al., 2011). Consequently, it is likely that records for biostratigraphically 580 important organisms have been over-inflated compared to other molluscs and between 581 localities. Koch (1978) showed by comparing previously existing collections and newly 582 collected records for the upper Cenomanian Sciponoceras gracile zone (now the Vascoceras 583 diartianum and Euomphaloceras septemseriatum zones; Cobban et al, 2006) that ammonites were better studied and more commonly reported than bivalve fauna. Parts of these collections 584 585 have made up the majority of the publicly available records of fossil occurrences within the 586 Western Interior, which are utilized in this study. As such, it is possible that ammonites are 587 over-represented in the early Campanian dataset and are skewing perceived results. However, 588 it is still possible to suggest that a suppressed expression of spatial aragonite bias occurs in the 589 distal offshore, albeit at reduced levels in comparison to the Cenomanian-Turonian interval.

590

591 Range size

Range size results reported a difference between calcitic and aragonitic bivalves across the twotime intervals studied, with aragonitic fauna showing a significantly smaller range size during

594 the Cenomanian-Turonian but not the Campanian. This variation is also expressed spatially 595 (Fig. 9). Within the Cenomanian-Turonian time slice, aragonitic geographic ranges (Fig. 9a) 596 are generally restricted to the western and northern edges of the seaway in comparison to 597 calcitic geographic ranges, which extend further to the center of the basin, as well as the east 598 and south (Fig. 9c). This same pattern is slightly different in the early Campanian interval (Fig. 599 9b,d); whilst aragonitic fauna still show a limited range, the difference between both bivalve 600 groups is less pronounced. This pattern also matches with the distribution of carbonate 601 deposition within the WIS: the Cenomanian-Turonian interval experienced widespread 602 carbonate sedimentation – in the form of the Greenhorn Limestone Formation – in the basin 603 center (Miall et al., 2008), whilst deposition in the basin center transitioned from limestones of 604 the Niobrara Formation to the siliciclastic mudstones of the Pierre Shale in the early Campanian 605 (McGookey et al., 1972; Da Gama et al., 2014). As our results confirm that carbonate 606 environments can exacerbate the effects of aragonite dissolution, it is possible that the 607 differences between the Cenomanian-Turonian and the Campanian are partially driven by the 608 enhanced effects of aragonite bias in carbonate-rich environments, resulting in a lowered sampling probability within carbonate-dominated localities. 609

610

# 611 Occurrence and Diversity Results

Overall, there is some evidence of aragonite dissolution influencing patterns of pure occurrences, taxonomic and subsampled diversity for aragonitic fauna, as previously hypothesized. In the Cenomanian-Turonian, aragonite bias is most pronounced within the proximal offshore bathymetric zone, with a lesser impact within the distal offshore zone. Whilst all fauna show a close correlation to collection counts for depth zones, both aragonitic and calcitic fauna deviate from this correlation in the proximal offshore zone, recording lower raw occurrences and diversity. The same is broadly observed in the Campanian: maximum

disparity of sampling probability between calcitic and aragonitic fauna is observed within the distal offshore zone, where aragonitic occurrences and raw taxic diversity show a noticeable decline and subsequent deviation from sampling proxies. Foote et al. (2015) reported similar results when comparing sampling-corrected results to ones that previously displayed the proportion of aragonitic taxa (Crampton et al., 2006), and concluded that similarities existed between sampling probabilities and relative proportions of aragonitic species.

Despite the potential relationships discussed above, we cannot report conclusive evidence for aragonite bias influencing the sampled diversity of molluscan fauna within the WIS. This aligns with other recent studies showing that despite evidence of widespread aragonite dissolution during early shallow diagenesis, perceived diversity is not largely affected by these processes (Behrensmeyer et al., 2005; Kidwell, 2005; Crampton et al., 2006; Hsieh et al., 2019). Hence, we must additionally look at external influences which might capture, enhance, or control the distribution of aragonitic faunas that would otherwise be lost to preferential dissolution.

632 Known human influences have potentially contributed to the suppression of aragonite bias on 633 a spatial scale. Whilst the extent to which aragonite dissolution may have influenced our 634 perceived record of molluscan diversity within the WIS is unclear, it is apparent that these 635 records closely correlate with established sampling proxies. Results of Spearmans-rank 636 correlation tests of occurrences and raw taxic diversity against sampling proxies for distance-637 from-paleoshoreline zones (Table 9) correlate strongly and significantly. It is clear that broader 638 scale sampling trends related to collector effort strongly influence the pattern of faunal 639 distribution across the seaway, potentially overwriting the effects of aragonite dissolution.

Whilst there have been many cases of preferential aragonite dissolution within local studies, aragonitic molluscan fauna are relatively well represented in the global fossil record (Harper, 1998). This paradox suggests that processes must occur which capture records of molluscan fauna at a higher frequency than they are capable of being destroyed. Cherns et al. (2008, 2011)

describe "Taphonomic Windows" as events in the fossil record which capture an unbiased view 644 645 of aragonitic faunas which have escaped preferential dissolution, and detail numerous 646 examples that may have operated within the WIS. One such window that is prevalent within 647 the WIS are concretions, sedimentary mineral masses of varying chemical composition that often form at shallow burial depths early in diagenesis when mineral cement precipitates 648 649 locally during lithification (Berner, 1968; McCoy et al., 2015). These have the potential to 650 preserve three-dimensional fossilized remains, often in exquisite detail (Dean et al., 2015; Korn 651 and Pagnac, 2017). Concretions are also a characteristic mode of molluscan occurrences within 652 the WIS, with fossil-bearing concretions found commonly throughout the seaway (Landman 653 and Klofak, 2012); as such, they could further contribute to a potential anthropogenic bias in 654 that they provide easily spotted locations to find fauna in otherwise barren strata (such as the 655 Pierre Shale), skewing collection intensity between localities with concretions and those without. However, only ~3% of USGS collections were obtained by selective collecting (Koch, 656 657 1980), and as USGS records make up ~55% of our finished dataset this suggests that sampling 658 intensity bias might be partially mitigated. Sediment accumulation rate could exert a large 659 influence on the potential for preferential aragonite dissolution to affect spatial zones of the sea 660 floor. If sediment accumulation rates were low, fauna would remain within the TAZ for an 661 extended period of time, and thus are more likely to be removed through physical reworking, 662 bioerosion and enhanced dissolution (Cherns et al., 2011). In contrast, if sediment 663 accumulation rates were high, fauna are likely to have been rapidly buried and thus have 664 escaped into the sub-TAZ region, where vulnerable bioclasts are likely to be stabilized by shallow burial diagenesis (Melim et al., 2002, 2004). Sediment accumulation rates within the 665 666 WIS varied both longitudinally within a stratigraphic interval (with higher sediment accumulation rates towards the western paleoshoreline) and with increased bathymetry in a 667 single location (Arthur and Sageman, 2005): accounting for this potential influence is 668

problematic, and the extent of its effects is ambiguous. The result of these factors is a potential
suppression of the spatial influence of aragonite dissolution bias on recorded faunal diversity
within the WIS.

672

673 Spatial Scale and Influence of Bias

674 The issue of scale is key to understanding the spatial impact of aragonite dissolution (Kosnik 675 et al., 2011). Foote et al. (2015) recorded preferential aragonite bias within carbonate-rich 676 environments on the regional spatial ( $\sim 10^6 \text{ km}^2$ ) and stage-level temporal (1-10 Myr) scales. 677 However, others (Behrensmeyer et al, 2005; Kidwell, 2005; Kiessling et al., 2008; Kosnik et 678 al, 2011) using global-scale data have reported negligible influence of shell mineralogy on 679 temporal trends or frequency of occurrences. Foote et al. (2015) reported three key differences 680 between previous studies and their work: higher taxonomic level of occurrences, larger time 681 bins, and the use of global data. These factors were inferred to "even out" spatial and temporal 682 variations in sampling, mitigating the influence and effect of locally variable biases inherent to 683 the fossil record. Foote et al. (2015) further suggested that as their taxonomic and temporal 684 scales were consistent with previously published work, an increase in spatial scale may prove 685 the most influential factor on demoting the influence of aragonite dissolution.

686 This result can be easily translated into the spatial expression of aragonite bias by comparing 687 its potential on alpha (within-site), beta (between-site) and gamma (global) diversity. At the 688 alpha level, the impact of aragonite bias on a single species will be at its most severe, 689 particularly within single-bed assemblages (Wright et al, 2003; Bush and Bambach, 2004; 690 Cherns et al 2008, 2011). However, at gamma levels of diversity, the probability of not 691 recording an individual drops substantially due to the number of possible localities to sample 692 from, where various taphonomic windows may result in aragonite preservation. As such, an 693 increased number of localities in a spatial setting are likely to partially obscure localized

aragonite dissolution. As we recorded an impact on zoned sampling probabilities and range size of aragonitic fauna in the WIS, but could not conclusively prove an influence on total diversity estimates, our data support the suggestions of Foote et al. (2015) that spatial scale is a dominant factor on the severity of aragonite bias.

698 Whilst unlikely to influence diversity on a global scale, this study has shown that preferential 699 aragonite dissolution has the capacity to govern the sampling probability of a species in 700 geographic space, and thus can influence the 'variation' definition of beta diversity (Anderson 701 et al., 2011). As the preferential dissolution of aragonite is a process that is exacerbated by 702 certain environments (Foote et al. 2015), its influence will impact localities with different 703 environmental conditions to differing extents – a species will be lost at one site and recorded 704 at another. Our results confirm this, showing aragonite bias has an effect on observed diversity 705 between locations, at least during times of widespread carbonate deposition.

706 Consequently, when looking at the spatial signal of aragonite dissolution as a whole, we can 707 see a sliding scale of influence: strong, environmentally dependent impact on alpha diversity; 708 a potentially large influence on beta diversity; and a negligible impact on gamma diversity. 709 Bush et al. (2004) grouped biases affecting spatially organized biodiversity in similar alpha, 710 beta and gamma levels, with alpha biases influencing within site diversity and beta and gamma 711 arising from failure to sample all available habitats or environments within a region. Whilst it 712 was noted in this study that taphonomic effects were not included in this definition, this system 713 can be modified in the light of our results. Aragonite bias, whilst operating at an alpha bias 714 (local) level, evidently has the capacity to systematically influence estimates of beta diversity. 715 As such, the influence of some taphonomic biases may be dependent on the spatial scale at 716 which they are observed. This is an important consideration for studies of the spatial 717 distribution of bias in the fossil record (Barnosky et al., 2005; Vilhena and Smith, 2013; Benson 718 et al., 2016; Close et al., 2017), and for paleobiogeographic studies in general.

719

# 720 CONCLUSIONS

721 1) A multifaceted approach shows that preferential aragonite dissolution is spatially 722 variable and impacts on the relative likelihood, absolute sampling probabilities, and 723 range sizes of aragonitic organisms within the Cretaceous Western Interior Seaway of 724 North America for a time interval that straddles the Cenomanian-Turonian boundary. 725 A similar but reduced effect is additionally observed within an early Campanian time 726 interval. A combination of depositional lithology (a limestone dominated basin within 727 the Cenomanian-Turonian compared to a more siliciclastic setting in the early 728 Campanian) and an anoxic basin center are hypothesized as drivers for this effect.

2) Carbonate environments enhance the effects of aragonite dissolution and the
preservation of calcitic organisms, as has been previously demonstrated. However, in
contrast to previous studies, siliciclastic environments are also shown to be influenced
by preferential aragonite dissolution.

3) Whilst similarities are observed between faunal distribution and absolute sampling
probabilities, we cannot conclusively say that aragonite dissolution has influenced
perceived diversity of molluscs within the Western Interior Seaway. "Taphonomic
windows" act to preserve records of organisms that would otherwise be lost. Other
anthropogenic and geologic biases appear to have a more obvious effect on the
molluscan record, and likely mask the influence of aragonite dissolution.

Whilst aragonite bias can be thought of as an "alpha bias", results show it could have a
systematic and severe impact on beta diversity. This suggests that taphonomic biases
can act differently at different scales in the spatial realm.

742

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| 982        | Figure and Table Captions   |
| 983        |   |
| 984        | Fig. 1.   |
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| 986        | Diagram showing potential model of spatial aragonite bias within the WIS. Within the outer shelf,       |
| 987        | preferential dissolution of aragonitic fauna is common, which has the potential to be expressed         |
| 988        | spatially. Within the basin center, anoxia limits benthic organism development, but allows for          |
| 989        | preservation of aragonitic material. Modified after Cherns et al. (2011).                               |
| 990        |   |
| 991        | Fig. 2.   |
| 992        |   |
| 993        | Paleogeographic zoned maps of the WIS used in this study. Depth-based zones are designated as           |
| 994        | nearshore, proximal offshore, distal offshore and basin center (Fig. 1); A. Paleobathymetric map of     |
| 995        | the Cenomanian-Turonian; B. Paleobathymetric map of the early Campanian.                                |
|            |   |
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- 996
- 997 Fig. 3. 998

Diagram showing potential model of apparent range size reduction due to spatially variable aragonite
preservation. Assuming that calcitic and aragonitic species of bivalve were both living at four separate
localities but aragonitic dissolution strongly influenced one of those locations (A), the resulting
convex hull for the aragonitic fauna drawn from surviving fossil occurrences would likely be smaller
than that of the calcitic organism (B).

- 1005
- 1005 Fig. 4.
- 1006

1007 Plots of generic level M2 sampling probabilities for the Cenomanian-Turonian (A, C, E, G) and lower 1008 Campanian (B, D, F, H) time slices across depth zones, split into carbonate and siliciclastic sampling 1009 opportunities. All results are plotted with percentage of carbonate collections per depth zone A. 1010 Cenomanian-Turonian generic level sampling probability, plotted with percentage of carbonate 1011 collections per depth zone; B. lower Campanian generic level sampling probability, plotted with 1012 percentage of carbonate collections per depth zone; C. Cenomanian-Turonian ammonite sampling 1013 probability; D. lower Campanian ammonite sampling probability; E. Cenomanian-Turonian aragonitic 1014 bivalve sampling probability; F. lower Campanian aragonitic bivalve sampling probability; E.

1015 Cenomanian-Turonian calcitic bivalve sampling probability; F. lower Campanian calcitic bivalve sampling probability.

- 1017
- 1018 Fig. 5
- 1019

Graph summarizing multiple logistic regression model results (T1-T8 = Tables 1-8). Final models are presented within columns, whereas factors are presented along rows for those models. Results are presented as either green (comparatively positive odds of sampling compared to reference factor), red (comparatively negative odds of sampling compared to reference factor), light grey (factor not statistically significant) and dark grey (factor not included in the final model). The following factors are used as a baseline for comparison: Mineralogy = aragonite; lithology = mudstone, lithology with mineralogy = aragonite:mudstone; zone = Basin Center (BC); diet = carnivore. Note that the

1027 magnitude of regression coefficients is not presented within this graph. NS: Near Shore; POS:

- 1028 Proximal Offshore; DOS: Distal offshore; BC: Basin Center.
- 1029
- 1030 Fig. 6. 1031

Range size plots for the Cenomanian-Turonian and lower Campanian. A. Cenomanian-Turonian box plots of range size for both aragonitic bivalves and calcitic bialves on log scale; B. lower Campanian box plots of range size for both aragonitic bivalves and calcitic bivalves on log scale; C. Randomized bootstrap for Cenomanian-Turonian mean range sizes – recorded difference in the mean is shown to be statistically significant; D. Randomized bootstrap for lower Campanian mean range sizes – recorded difference in the mean is not shown to be statistically significant.

1038

1039 Fig. 7. 1040

Plots of generic level diversity plots for the Cenomanian-Turonian and lower Campanian within depth
zones, plotted with number of collections and outcrop area. A. Generic diversity and number of
collections for the Cenomanian-Turonian; B. Generic diversity and number of collections for the
lower Campanian; C. Generic diversity and outcrop area for the Cenomanian-Turonian; D. Generic
diversity and outcrop area for the lower Campanian.

1046 1047 Fig. 8.

1048

Plots of generic level SQS results for depth zones in the Cenomanian-Turonian and lower Campanian,
set at 0.4, 0.5 and 0.6 quora. A. SQS results for ammonites in the Cenomanian-Turonian; B. SQS

| 1051<br>1052<br>1053<br>1054<br>1055 | results for ammonites in the lower Campanian; C. SQS results for aragonitic bivalves in the Cenomanian-Turonian; D. SQS results for aragonitic bivalves in the lower Campanian; E. SQS results for calcitic bivalves in the Cenomanian-Turonian; F. SQS results for calcitic bivalves in the lower Campanian.  |
|--------------------------------------|--|
| 1055<br>1056<br>1057                 | Fig. 9.  |
| 1059<br>1059<br>1060<br>1061         | Paleogeographic maps shown with range sizes of calcitic and aragonitic bivalves for both time slices.<br>A. Aragonitic bivalve range sizes for the Cenomanian-Turonian; B. Aragonitic bivalve range sizes for the lower Campanian; C. Calcitic bivalve range sizes for the Cenomanian-Turonian; D. Calcitic bivalve range sizes for the lower Campanian. |
| 1062<br>1063<br>1064                 | Table 1.   |
| 1065<br>1066<br>1067                 | Table for Multiple Logistic Regression results for all bivalves within the Cenomanian-Turonian across the whole seaway, using model with lowest AIC score. M - Mineralogy; D - Diet; L - Lithology; Z – depth zone.  |
| 1068<br>1069                         | Table 2.   |
| 1070<br>1071                         | Table for Multiple Logistic Regression results for all bivalves within the Cenomanian-Turonian for each depth zone, using models with lowest AIC scores. M - Mineralogy; D - Diet; L – Lithology.  |
| 1072<br>1073                         | Table 3.   |
| 1074<br>1075<br>1076                 | Table for Multiple Logistic Regression results for all organisms (including ammonites) within the Cenomanian-Turonian across the whole seaway, using model with lowest AIC score. M - Mineralogy; L - Lithology; Z – depth zone.   |
| 1077<br>1078<br>1079                 | Table 4.   |
| 1080<br>1081<br>1082<br>1083         | Table for Multiple Logistic Regression results for all organisms (including ammonites) within the Cenomanian-Turonian for each depth zone, using models with lowest AIC scores. M - Mineralogy; L – Lithology.   |
| 1085<br>1084<br>1085                 | Table 5.   |
| 1086<br>1087<br>1088<br>1089         | Table for Multiple Logistic Regression results for all bivalves within the lower Campanian across the whole seaway, using model with lowest AIC score. M - Mineralogy; D - Diet; L - Lithology; Z – depth zone.  |
| 1089<br>1090<br>1091                 | Table 6.   |
| 1091<br>1092<br>1093<br>1094         | Table for Multiple Logistic Regression results for all bivalves within the lower Campanian for each depth zone, using models with lowest AIC scores. M - Mineralogy; D - Diet; L - Lithology; Z – depth zone.  |
| 1095<br>1096<br>1097                 | Table 7.   |
| 1098<br>1099<br>1100                 | Table for Multiple Logistic Regression results for all organisms (including ammonites) within the lower Campanian across the whole seaway, using model with lowest AIC score. M - Mineralogy; L - Lithology; Z – depth zone.   |
| 1101<br>1102<br>1103                 | Table 8.   |

- 1104 Table for Multiple Logistic Regression results for all organisms (including ammonites) within the lower
- 1105 Campanian for each depth zone, using models with lowest AIC scores. M Mineralogy; L Lithology.
- 1106 1107 Table 9.
- 1108
- 1109 Spearman's rank correlations between generic diversity of faunal groups and various sampling proxies
- 1110 for distance-from-paleoshoreline zones within the Cenomanian-Turonian and lower Campanian.

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Figure 1. Diagram showing potential model of spatial aragonite bias within the WIS. Within the outer shelf, preferential dissolution of aragonitic fauna is common, which has the potential to be expressed spatially. Within the basin center, anoxia limits benthic organism development, but allows for preservation of aragonitic material. Modified after Cherns et al. (2011).



Figure 2. Paleogeographic zoned maps of the WIS used in this study. Depth-based zones are designated as nearshore, proximal offshore, distal offshore and basin center (Fig. 1); A. Paleobathymetric map of the Cenomanian-Turonian; B. Paleobathymetric map of the early Campanian.



Figure 3. Diagram showing potential model of apparent range size reduction due to spatially variable aragonite preservation. Assuming that calcitic and aragonitic species of bivalve were both living at four separate localities but aragonitic dissolution strongly influenced one of those locations (A), the resulting convex hull for the aragonitic fauna drawn from surviving fossil occurrences would likely be smaller than that of the calcitic organism (B).



Plots of generic level M2 sampling probabilities for the Cenomanian-Turonian (A, C, E, G) and lower Campanian (B, D, F, H) time slices across depth zones, split into carbonate and siliciclastic sampling opportunities. All results are plotted with percentage of carbonate collections per depth zone A.
 Cenomanian-Turonian generic level sampling probability, plotted with percentage of carbonate collections per depth zone; B. lower Campanian generic level sampling probability, plotted with percentage of carbonate collections per depth zone; C. Cenomanian-Turonian ammonite sampling probability; D. lower Campanian ammonite sampling probability; E. Cenomanian-Turonian aragonitic bivalve sampling probability; F. lower Campanian aragonitic bivalve sampling probability; F. lower Campanian calcitic bivalve sampling probability; F. lower Campanian calcitic bivalve sampling probability.

| Comparatively positive odds   |   | CENOMANIAN-TURONIAN |    |     |               |    | CAMPANIAN |    |          |     |    |               |    |     |     |    |           |    |     |     |    |
|-------------------------------|---|---------------------|----|-----|---------------|----|-----------|----|----------|-----|----|---------------|----|-----|-----|----|-----------|----|-----|-----|----|
| Comparatively negative odds   |   | Bivalves            |    |     | All Organisms |    |           |    | Bivalves |     |    | All Organisms |    |     |     |    |           |    |     |     |    |
| Facto                         | or not statistically significant  | T1                  |    | T2  | 2             |    | Т3        |    | T4       | 1   |    | T5            |    | T   | 6   |    | T7        |    | Τŧ  | }   |    |
| Facto                         | or not included in final model  | All basin           | NS | POS | DOS           | BC | All basin | NS | POS      | DOS | BC | All basin     | NS | POS | DOS | BC | All basin | NS | POS | DOS | рС |
| Mineralogy                    | Calcite   |                     |    |     |               |    |           |    |          |     |    |               |    |     |     |    |           |    |     |     |    |
| Lithology                     | Calcarenite<br>Calcareous shale<br>Chalk<br>Conglomerate<br>Limestone<br>Marl<br>Sandstone<br>Sittstone   |                     |    |     |               |    |           |    |          |     |    |               | 1  |     |     |    |           |    |     |     |    |
| Lithology with<br>Mineralogy: | Calcite:Calcarenite<br>Calcite:Calcareous shale<br>Calcite:Calcareous shale<br>Calcite:Chalk<br>Calcite:Conglomerate<br>Calcite:Limestone<br>Calcite:Marl<br>Calcite:Sandstone<br>Calcite:Siltstone |                     |    |     |               |    |           |    |          |     |    |               |    |     |     |    |           |    |     |     |    |
| Zone                          | NS<br>POS<br>DOS  |                     |    |     |               |    |           |    |          |     |    |               |    |     |     |    |           |    |     |     |    |
| Diet                          | Suspension feeder<br>Chemosymbiont deposit feeder<br>Deposit feeder<br>Subsurface deposit feeder<br>Herbivore   |                     |    |     |               |    |           |    |          |     |    |               |    |     |     |    |           |    |     |     |    |

Graph summarizing multiple logistic regression model results (T1-T8 = Tables 1-8). Final models are presented within columns, whereas factors are presented along rows for those models. Results are presented as either green (comparatively positive odds of sampling compared to reference factor), red (comparatively negative odds of sampling compared to reference factor), light grey (factor not statistically significant) and dark grey (factor not included in the final model). The following factors are used as a baseline for comparison: Mineralogy = aragonite; lithology = mudstone, lithology with mineralogy = aragonite:mudstone; zone = Basin Center (BC); diet = carnivore. Note that the magnitude of regression coefficients is not presented within this graph. NS: Near Shore; POS: Proximal Offshore; DOS: Distal offshore; BC: Basin Center.



Range size plots for the Cenomanian-Turonian and lower Campanian. A. Cenomanian-Turonian box plots of range size for both aragonitic bivalves and calcitic bialves on log scale; B. lower Campanian box plots of range size for both aragonitic bivalves and calcitic bivalves on log scale; C. Randomized bootstrap for Cenomanian-Turonian mean range sizes – recorded difference in the mean is shown to be statistically significant; D. Randomized bootstrap for lower Campanian mean range sizes – recorded difference in the mean is not shown to be statistically significant.



Plots of generic level diversity plots for the Cenomanian-Turonian and lower Campanian within depth zones, plotted with number of collections and outcrop area. A. Generic diversity and number of collections for the Cenomanian-Turonian; B. Generic diversity and number of collections for the lower Campanian; C. Generic diversity and outcrop area for the Cenomanian-Turonian; D. Generic diversity and outcrop area for the lower Campanian.



Plots of generic level SQS results for depth zones in the Cenomanian-Turonian and lower Campanian, set at 0.4, 0.5 and 0.6 quora. A. SQS results for ammonites in the Cenomanian-Turonian; B. SQS results for ammonites in the lower Campanian; C. SQS results for aragonitic bivalves in the Cenomanian-Turonian; D. SQS results for aragonitic bivalves in the lower Campanian; F. SQS results for calcitic bivalves in the lower Campanian.



Paleogeographic maps shown with range sizes of calcitic and aragonitic bivalves for both time slices. A. Aragonitic bivalve range sizes for the Cenomanian-Turonian; B. Aragonitic bivalve range sizes for the lower Campanian; C. Calcitic bivalve range sizes for the Cenomanian-Turonian; D. Calcitic bivalve range sizes for the lower Campanian.

| Final Model        | AIC      | Factor                         | Regression coefficient | Standard error |
|--------------------|----------|--------------------------------|------------------------|----------------|
| M + D + L + Z + M: | L 22367* | M-Calcite                      | 1.516                  | 0.080          |
|                    |          | D-Chemosymbiont deposit feeder | 1.353                  | 0.239          |
|                    |          | L-Sandstone                    | 0.672                  | 0.115          |
|                    |          | L-Silt                         | 0.759                  | 0.362          |
|                    |          | M-Calcite:L-Limestone          | 1.338                  | 0.192          |
|                    |          | L-Limestone                    | -1.415                 | 0.184          |
|                    |          | DOS                            | -0.410                 | 0.082          |
|                    |          | NS                             | -0.754                 | 0.114          |
|                    |          | POS                            | -0.684                 | 0.089          |
|                    |          | D-Herbivore                    | -1.870                 | 1.022          |
|                    |          | M-Calcite:L-Calcarenite        | 1.146                  | 0.611          |

Table for Multiple Logistic Regression results for all bivalves within the Cenomanian-Turonian across the who

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| p-value      |
|--------------|
| < 2e-16 ***  |
| 1.45e-08 *** |
| 5.39e-09 *** |
| 0.0359 *     |
| 3.35e-12 *** |
| 1.49e-14 *** |
| 6.70e-07 *** |
| 4.30e-11 *** |
| 1.10e-14 *** |
| 0.0673 .     |
| 0.0605 .     |

ble seaway, using model with lowest AIC score. M - Mineralogy; D - Diet; L - Lithology; Z – depth zone.

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| Data              | Final Model     | AIC    | Factor                         |
|-------------------|-----------------|--------|--------------------------------|
| Noar Shoro        | M + L           | 1808.7 | M-Calcite                      |
| Near Shore        |                 |        | L-Sandstone                    |
|                   | M + D + L + M:L | 7809.2 | M-Calcite                      |
|                   |                 |        | D-Chemosymbiont deposit feeder |
| Drovimal Offshoro |                 |        | L-Marl                         |
|                   |                 |        | M-Calcite:L-Limestone          |
|                   |                 |        | L-Limestone                    |
|                   |                 |        | M-Calcite:L-Marl               |
|                   | M + L + D + M:L | 10768  | M-Calcite                      |
|                   |                 |        | L-Sandstone                    |
| Dictal Offchara   |                 |        | L-Silt                         |
| Distal Offshore   |                 |        | D-Chemosymbiont deposit feeder |
|                   |                 |        | M-Calcite:L-Limestone          |
|                   |                 |        | L-Limestone                    |
|                   | M + L + D       | 1857.1 | L-Calcarenite                  |
| Basin Center      |                 |        | L-Calcareous Shale             |
|                   |                 |        | L-Marl                         |

Table for Multiple Logistic Regression results for all bivalves within the Cenomanian-Turoni

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| Regression coefficient | Standard error | p-value      |
|------------------------|----------------|--------------|
| 0.626                  | 0.176          | 0.000372 *** |
| 0.571                  | 0.166          | 0.000593 *** |
| 1.818                  | 0.134          | < 2e-16 ***  |
| 1.917                  | 0.449          | 1.94e-05 *** |
| 3.125                  | 0.539          | 6.89e-09 *** |
| 1.390                  | 0.438          | 0.00149 **   |
| -1.358                 | 0.422          | 0.00131 **   |
| -1.775                 | 0.808          | 0.02804 *    |
| 1.416                  | 0.111          | < 2e-16 ***  |
| 0.978                  | 0.152          | 1.18e-10 *** |
| 1.684                  | 0.595          | 0.004671 **  |
| 1.051                  | 0.293          | 0.000335 *** |
| 1.222                  | 0.226          | 6.08e-08 *** |
| -1.248                 | 0.211          | 3.25e-09 *** |
| 1.777                  | 0.582          | 0.00225 **   |
| 1.826                  | 0.371          | 8.47e-07 *** |
| 1.864                  | 0.489          | 0.00014 ***  |

ian for each depth zone, using models with lowest AIC scores. M - Mineralogy; D - Diet; L – Lithology.

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| Final Model     | AIC    | Factor                       | Regression coefficient | Standard error |
|-----------------|--------|------------------------------|------------------------|----------------|
| M + Z + L + M:L | 46464* | M-Calcite                    | 0.741                  | 0.044          |
|                 |        | L-Conglomerate               | 1.495                  | 0.324          |
|                 |        | L-Marl                       | 0.593                  | 0.174          |
|                 |        | M-Calcite:L-Calcarenite      | 1.269                  | 0.319          |
|                 |        | M-Calcite:L-Calcareous Shale | 2.290                  | 0.280          |
|                 |        | M-Calcite:L-Limestone        | 0.202                  | 0.072          |
|                 |        | M-Calcite:L-Sandstone        | 0.595                  | 0.095          |
|                 |        | DOS                          | -0.530                 | 0.050          |
|                 |        | NS                           | -0.700                 | 0.076          |
|                 |        | POS                          | -0.755                 | 0.056          |
|                 |        | L-Calcarenite                | -0.959                 | 0.260          |
|                 |        | L-Calcareous Shale           | -1.561                 | 0.253          |
|                 |        | L-Limestone                  | -0.288                 | 0.050          |
|                 |        | L-Sandstone                  | -0.152                 | 0.067          |
|                 |        | M-Calcite:L-Conglomerate     | -1.857                 | 1.057          |

Table for Multiple Logistic Regression for all organisms (including ammonites) within the Cenomanian-Ture

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| p-value      |
|--------------|
| < 2e-16 ***  |
| 3.92e-06 *** |
| 0.000654 *** |
| 7.10e-05 *** |
| 3.03e-16 *** |
| 0.004996 **  |
| 4.31e-10 *** |
| < 2e-16 ***  |
| < 2e-16 ***  |
| < 2e-16 ***  |
| 0.000227 *** |
| 6.77e-10 *** |
| 6.82e-09 *** |
| 0.024007 *   |
| 0.079016 .   |

onian across the whole seaway, using model with lowest AIC score. M - Mineralogy; L - Lithology; Z - depth zc

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| Data              | Final Model | AIC     | Factor                       | Regression coefficient |
|-------------------|-------------|---------|------------------------------|------------------------|
| Near Shore        | М           | 3563.3  | M-Calcite                    | 0.443                  |
|                   | M + L + M:L | 14954*  | M-Calcite                    | 0.942                  |
|                   |             |         | L-Conglomerate               | 1.553                  |
|                   |             |         | L-Marl                       | 2.046                  |
|                   |             |         | M-Calcite:L-Calcareous Shale | 1.900                  |
| Proximal Offshore |             |         | M-Calcite:L-Sandstone        | 0.368                  |
|                   |             |         | L-Calcareous Shale           | -1.765                 |
|                   |             |         | L-Sandstone                  | -0.261                 |
|                   |             |         | L-Limestone                  | -0.203                 |
|                   |             |         | M-Calcite:L-Conglomerate     | -2.057                 |
|                   | M + L + M:L | 21160*  | M-Calcite                    | 0.701                  |
|                   |             |         | L-Marl                       | 0.822                  |
|                   |             |         | M-Calcite:L-Calcarenite      | 1.639                  |
|                   |             |         | M-Calcite:L-Calcareous Shale | 1.842                  |
|                   |             |         | M-Calcite:L-Limestone        | 0.287                  |
| Distal Offshore   | (           |         | M-Calcite:L-Sandstone        | 0.849                  |
|                   |             |         | L-Calcarenite                | -1.530                 |
|                   |             |         | L-Calcareous Shale           | -1.651                 |
|                   |             |         | L-Limestone                  | -0.313                 |
|                   |             |         | M-Calcite:L-Marl             | -1.747                 |
|                   |             |         | M-Calcite:L-Silt             | 1.074                  |
|                   | M + L + M:L | 6692.3* | L-Marl                       | -2.270                 |
|                   |             |         | M-Calcite:L-Calcarenite      | 3.536                  |
| Basin Center      |             |         | M-Calcite:L-Calcareous Shale | 3.479                  |
|                   |             |         | M-Calcite:L-Limestone        | 0.868                  |
|                   |             |         | M-Calcite:L-Marl             | 4.135                  |
|                   |             |         |                              |                        |

Table for Multiple Logistic Regression results for all organisms (including ammonites) within the Cenomania

| Standard error | p-value      |
|----------------|--------------|
| 0.114          | 9.92e-05 *** |
| 0.070          | < 2e-16 ***  |
| 0.325          | 1.79e-06 *** |
| 0.367          | 2.47e-08 *** |
| 0.636          | 0.00281 **   |
| 0.153          | 0.01635 *    |
| 0.578          | 0.00224 **   |
| 0.108          | 0.01584 *    |
| 0.108          | 0.06071.     |
| 1.059          | 0.05197 .    |
| 0.064          | < 2e-16 ***  |
| 0.235          | 0.000475 *** |
| 0.583          | 0.004940 **  |
| 0.657          | 0.005042 **  |
| 0.105          | 0.006111 **  |
| 0.141          | 1.60e-09 *** |
| 0.502          | 0.002308 **  |
| 0.579          | 0.004370 **  |
| 0.067          | 3.15e-06 *** |
| 0.750          | 0.019823 *   |
| 0.588          | 0.067947 .   |
| 1.011          | 0.024660 *   |
| 1.167          | 0.002442 **  |
| 0.507          | 6.53e-12 *** |
| 0.376          | 0.021101 *   |
| 1.123          | 0.000231 *** |

an-Turonian for each depth zone, using models with lowest AIC scores. M - Mineralogy; L – Litho

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| Final Model | AIC     | Factor                | Regression coefficient | Standard error |
|-------------|---------|-----------------------|------------------------|----------------|
| M + L + M:L | 7063.9* | M-Calcite             | 0.767                  | 0.115          |
|             |         | L-Sandstone           | 0.594                  | 0.168          |
|             |         | M-Calcite:L-Limestone | 1.292                  | 0.538          |
|             |         | M-Calcite:L-Sandstone | -0.496                 | 0.209          |
|             |         | M-Calcite:L-Siltstone | 1.838                  | 1.044          |

Table for Multiple Logistic Regression results for all bivalves within the lower Campanian across the v

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| p-value      |
|--------------|
| 2.41e-11 *** |
| 0.00041 ***  |
| 0.01625 *    |
| 0.01781 *    |
| 0.07825 .    |

whole seaway, using model with lowest AIC score. M - Mineralogy; D - Diet; L - Lithology; Z – depth zone.

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| Data              | Final Model | AIC     | Factor                | Regression coefficient |
|-------------------|-------------|---------|-----------------------|------------------------|
| Near Shore        | L           | 886.47  | N/A                   | N/A                    |
|                   | M + L       | 1603.9  | M-Calcite             | 0.808                  |
| Proximal Offshore |             |         | L-Calcareous Shale    | 0.994                  |
|                   |             |         | L-Limestone           | 0.687                  |
|                   | M + L + M:L | 3420.6* | M-Calcite             | 1.145                  |
| Distal Offshore   |             |         | L-Sandstone           | 0.765                  |
|                   |             |         | M-Calcite:L-Sandstone | -0.709                 |
| Basin Center      | L+D         | 1158.2  | L-Calcareous Shale    | 0.812                  |

Table for Multiple Logistic Regression results for all bivalves within the lower Campanian for each d

| r p-value      | Standard error |
|----------------|----------------|
| N/A            | N/A            |
| 1.08e-05 ***   | 0.184          |
| ′ 0.03336 *    | 0.467          |
| ′ 0.00545 **   | 0.247          |
| ) 1.41e-11 *** | 0.169          |
| 0.00332 **     | 0.260          |
| 0.02768 *      | 0.322          |
| 0.000259 ***   | 0.222          |

lepth zone, using models with lowest AIC scores. M - Mineralogy; D - Diet; L - Lithology; Z - depth

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| Final Model | AIC   | Factor    | Regression coefficient | Standard error | p-value      |
|-------------|-------|-----------|------------------------|----------------|--------------|
| Z + M       | 16712 | DOS       | -0.323                 | 0.067          | 1.61e-06 *** |
|             |       | NS        | -0.203                 | 0.095          | 0.033311 *   |
|             |       | POS       | -0.281                 | 0.078          | 0.000307 *** |
|             |       | M-Calcite | -0.146                 | 0.054          | 0.006599 **  |

Table for Multiple Logistic Regression results for all organisms (including ammonites) within the lower Car

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mpanian across the whole seaway, using model with lowest AIC score. M - Mineralogy; L - Lithology; Z - dep

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th zone.

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| Data              | Final Model | AIC     | Factor    | Regression coefficient | Standard error |
|-------------------|-------------|---------|-----------|------------------------|----------------|
| Near Shore        | Rcoll ~ 1   | 1747.1  | N/A       | N/A                    | N/A            |
| Proximal Offshore | М           | 3683    | M-Calcite | -0.307                 | 0.117          |
| Distal Offshore   | Rcoll ~ 1   | 8319.3  | N/A       | N/A                    | N/A            |
| Basin Center      | L + M + L:M | 1151.9* | N/A       | N/A                    | N/A            |

Table for Multiple Logistic Regression results for all organisms (including ammonites) within the lo

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| p-value    |
|------------|
| N/A        |
| 0.00838 ** |
| N/A        |
| N/A        |

wer Campanian for each depth zone, using models with lowest AIC scores. M - Mineralogy; L – Lith

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|                   | DISTANCE       | E            | Correlation                                       | Spearman's<br>Rho |
|-------------------|----------------|--------------|---|-------------------|
| NIAN              | AMMONITES      | Outcrop Area | Ammonite occurrences vs. Outcrop Area             | 0.475             |
|                   |                |              | Ammonite raw diversity vs. Outcrop Area           | 0.704             |
|                   |                | Collections  | Ammonite occurrences vs. Collections              | 0.885             |
|                   |                |              | Ammonite raw diversity vs. Collections            | 0.732             |
| MANIAN-TURG       | ARAG. BIVALVES | Outcrop Area | Aragonitic bivalve occurrences vs. Outcrop Area   | 0.634             |
|                   |                |              | Aragonitic bivalve raw diversity vs. Outcrop Area | 0.640             |
|                   |                | Collections  | Aragonitic bivalve occurrences vs. Collections    | 0.568             |
|                   |                |              | Aragonitic bivalve raw diversity vs. Collections  | 0.502             |
| CENOI             | CALC. BIVALVES | Outcrop Area | Calcitic bivalve occurrences vs. Outcrop Area     | 0.534             |
|                   |                |              | Calcitic bivalve raw diversity vs. Outcrop Area   | 0.678             |
|                   |                | Collections  | Calcitic bivalve occurrences vs. Collections      | 0.904             |
|                   |                |              | Calcitic bivalve raw diversity vs. Collections    | 0.738             |
|                   | AMMONITES      | Outcrop Area | Ammonite occurrences vs. Outcrop Area             | 0.221             |
|                   |                |              | Ammonite raw diversity vs. Outcrop Area           | 0.343             |
| IAN               |                | Collections  | Ammonite occurrences vs. Collections              | 0.970             |
|                   |                |              | Ammonite raw diversity vs. Collections            | 0.727             |
| <b>WER CAMPAN</b> | ARAG. BIVALVES | Outcrop Area | Aragonitic bivalve occurrences vs. Outcrop Area   | 0.473             |
|                   |                |              | Aragonitic bivalve raw diversity vs. Outcrop Area | 0.404             |
|                   |                | Collections  | Aragonitic bivalve occurrences vs. Collections    | 0.822             |
|                   |                |              | Aragonitic bivalve raw diversity vs. Collections  | 0.759             |
| ΓΟ                | CALC. BIVALVES | Outcrop Area | Calcitic bivalve occurrences vs. Outcrop Area     | 0.056             |
|                   |                |              | Calcitic bivalve raw diversity vs. Outcrop Area   | 0.322             |
|                   |                | Collections  | Calcitic bivalve occurrences vs. Collections      | 0.958             |
|                   |                |              | Calcitic bivalve raw diversity vs. Collections    | 0.879             |

Table 9: Spearman's rank correlations between generic diversity of faunal groups and various samp

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| p-value | Sig. |
|---------|------|
| 0.05569 | N/A  |
| 0.00163 | *    |
| 0.00000 | *    |
| 0.00084 | *    |
| 0.00624 | *    |
| 0.00569 | *    |
| 0.01740 | *    |
| 0.04011 | *    |
| 0.02918 | *    |
| 0.00277 | *    |
| 0.00000 | *    |
| 0.00071 | *    |
| 0.49070 | N/A  |
| 0.27623 | N/A  |
| 0.00000 | *    |
| 0.01000 | *    |
| 0.12070 | N/A  |
| 0.19248 | N/A  |
| 0.00104 | *    |
| 0.00418 | *    |
| 0.86898 | N/A  |
| 0.30701 | N/A  |
| 0.00000 | *    |
| 0.00017 | *    |

ling proxies for distance-from-paleoshoreline zones within the Cenomanian-Turonian and lower Cam

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